# 510(K) SUMMARY AS REQUIRED BY SECTION 807.92(c)

### 1.- SUBMITTER INFORMATION:

Company Name:

Sauffon Pharmaceuticals Ltd.

Address:

49-53 York Street

Twickenham Middlesex TW13LP

Phone:

020.8322.4200

Fax:

020 8891 2833

Contact Person:

Dr Christopher Smejkal

DATE SUMMARY PREPARED: 15th October 2013

**DEVICE NAME:** 

Trade Name:

Sauflon Clariti 1 Day (somofilcon A) Soft

(hydrophilic) Daily Disposable Contact Lens with UV

**Blocker and Handling Tint** 

Common Name:

Soft Contact Lens

Classification:

CLASS II (21 CFR 886.5925) CODE -LPL, MVN

SOFT (HYDROPHILIC) CONTACT LENS

**Duration of Contact:** 

A daily replacement lens for surface contact of the eye

and intended for daily removal and less than 24 hours

contact duration.

### 2.- SUBSTANTIAL ÉQUIVALENCE:

The sponsor considers the Sauffon Clariti 1 Day (somofilcon, A) Soft (hydrophilic) Daily Disposable Contact Lens with UV Blocker and Handling Tint to be substantially equivalent to the Saufton Clariti 1 Day (somofileon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV Blocker, cleared pursuant to K130331, Acuvue Trueye (Narafilcon A) Soft (Hydrophilic) Visibility Tinted Contact Lens for Daily Wear single use which has been approved pursuant to K073485, and Air Optix (Lotrafilcon B) Soft (Hydrophilic) Visibility Tinted Contact Lens for Daily Wear which has been approved pursuant to K033919/K073459.

### 3.- DESCRIPTION of the DEVICE:

The Saufion Ctariti 1 Day (somofileon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV Blocker and Handling Tint is available as a single vision, toric, multifocal and multifocal toric lens. The lens material (somofileon A) is a hydrophilic co-polymer of silicone containing monomers and hydrophilic monomers which is cross-linked with tetraethyleneglycol dimethacrylate. When hydrated the lens consists of 44.0% somofileon A and 56.0% water by weight of saline immersed in normal saline. A benzophenone UV absorbing monomer is used to block UV radiation. A handling tint is added to the composition of the lens to make the lens visible for handling.

The average transmittance characteristics are less than 5% in the UVB range of 280 to 315nm and less than 50% in the UVA range of 316-380nm

The Sauffon Clariti 1 Day (somofileon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV Blocker and Handling Tint is a hemispherical flexible shell, which covers the cornea and a portion of the adjacent sclera, with the following dimensions:

	Chord Diameter:	13.0mm to 18.0mm
•	Centre Thickness:	0.03mm to 0.70mm
•	Base Curve:	7.5mm to 9.50mm
•	Powers:	-30.00 DS to +30.00 DS
	Toric Cylinder options:	-0.75 to -9.75
•	Toric Axis options:	$10^{\circ}$ to $180^{\circ}$ (5° steps).

Multifocal Add:

Add power up to +4.00 labelled with indicative add strength to be read in conjunction with the fitting guide.

The physical/optical properties of the lenses are:

•	Refractive Index:	1.4008
•	%Transmittance @ 590nm:	98.30
	%Transmittance @ 280-315nm:	0.71
•	%Transmittance @ 316-380nm:	20.62
•	Surface Character:	Hydrophilic
•	Water Content:	56%
•	Oxygen Permeability (DK):	60 x 10 <sup>-11</sup> (cm <sup>2</sup> /sec) (ml O2/ml x mmHg) at 35°C (Fatt Method for determination of oxygen permeability).
•:	Specific Gravity:	1.17

### 4.- INDICATIONS FOR USE

Sauffon Clariti 1 Day (somofilcon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV blocker and Handling Tint is indicated for:

The SAUFLON CLARITI 1 DAY (somofileon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV blocker and Handling Tint is indicated for daily wear single use only for the correction of refractive ametropia (myopia and hyperopia) in phakic or aphakic persons with non-diseased eyes that may exhibit astigmatism up to 2.00 Diopters that does not interfere with visual acuity.

The SAUFLON CLARITI I DAY TORIC (somofileon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV blocker and Handling Tint is indicated for daily wear single use only for the optical correction of refractive ametropia (myopia and hyperopia) in phasic or aphasic persons with non-diseased eyes that may exhibit astigmatism up to 10.00 Diopters.

The SAUFLON CLARITI 1 DAY MULTIFOCAL (somofileon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV blocker and Handling Tint is indicated for daily wear single use only for the optical correction of refractive ametropia (myopia and hyperopia) and/or presbyopia in phakic or aphakic persons with non-diseased eyes that may require a reading addition of +4.00 Diopters or less and may exhibit astigmatism up to 1.50 Diopters or less.

The SAUFLON CLARITI 1 DAY MULTIFOCAL TORIC (somofilcon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV blocker and Handling Tint is indicated for daily wear single use only for the optical correction of refractive ametropia (myopia and hyperopia) and/or presbyopia in phakic or aphakic persons with non-diseased eyes that may exhibit astigmatism up to 10.00 Diopters and require a reading addition of +4.00 Diopters or less.

Sauffon Clariti 1 Day (somofilcon A) Soft (hydrophilic) Daily Disposable Contact lens with UV blocker and Handling Tint help protect against transmission of harmful UV radiation to the cornea and into the eye.

### 5.- PREDICATE DEVICES

The sponsor considers the SAUFLON CLARITI IDAY (somofileon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV Blocker and Handling Tint to be substantially equivalent to the SAUFLON CLARITI 1DAY (somofileon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV Blocker cleared pursuant to K130331, Acuvue Trueye (Narafileon A) Soft (hydrophilic) Visibility Tinted Contact Lens for Daily Wear single use which has been approved pursuant to K073485, and Air Optix (Lotrafileon B) Soft (hydrophilic) Visibility Tinted Contact Lens for Daily Wear which has been approved pursuant to K033919/K073459.

The following table summarises the primary features for this comparison, illustrating the similarities and differences.

Comparison of Phy Blocker and handli	sical / Optical Properties for the Sing tint versus Acuvue Trueye (Narafil	Comparison of Physical / Optical Properties for the SAUFLON CLARITI 1 DAY (somofilcon A) Silicone Hydrogel Blocker and handling tint versus Acuvue Trueye (Narafilcon A) and SAUFLON CLARITI 1 DAY SILICONE HYDROGEL	Comparison of Physical / Optical Properties for the SAUFLON CLARITI 1 DAY (somofilcon A) Silicone Hydrogel Contact Lens with UV Blocker and handling tint versus Acuvue Trueye (Narafilcon A) and SAUFLON CLARITI 1 DAY SILICONE HYDROGEL
	PREDICATE DEVICE ACUVUE TRUEVE (K073485)	PREDICATE DEVICE - SAUFLON:CLARITI 1 DAY-SILICONE HYDROGEL (K130331)	SUBJECT DEVICE - SAUFLON CLARITI I DAY WITH HANDLING TINT
LENS MATERIAL	narafilcon A Silicone Hydrogel	symofileon A Silicone Hydrogel	somofileon A Silicone Hydrogel
INDICATIONS FOR Daily wear single use USE	Daily wear single use	Daily wear single use	Daily wear single use.
MANUFACTURING PROCESS	Cast Moulding.	Cast Moulding	Cast Moulding
WATER-CONTENT	47%	9696	969%
REFRACTIVE INDEX	1,41	1:40	1.40
LIGHT TRANSMITTANCE	85% minimum	\$696≤	%96₹
DK @35°C (RDGE CORRECTED)	100 (polarôgraphic method)	60 (Coulometric niethod)	60 (polarographic method)
COLOUR	Blue Visibility Tint	No Visibility Tim	Blue Visibility Tint

'I'NI'	CI Reactive Blue Dye 4	попс	D&C Green No. 6
UVBLOCKER	Benziariazole	Вспхорнепопе	Benzaphénone
MODULUS (MPa)	0,66	0.55	0.55
TIENSILE STRENGTH (MPa).	0,72	1.05	1.05
ELONGATION AT BREAK %	170	163	163
PĄCKAGING MĄTIBRJALS	Injected molded polypropylene blisters covered by aluminium foil luminate and blister strips are packed into printed cartons	Injected molded polypropylene blisters covered by aluminium foil laminate and blister strips are packed into printed cartons	Injected molded polypropylene blisters covered by aluminium foil faminate and blister strips are packed into printed cartons
PACKAGING SOLUTION	Buffered saline solution containing up to 0.01% methyl ether cellulose	Borate buffered saline solution containing 0.005% poloxamer	Borate buffered saline solution containing 0.005% poloxamer
PACKAGING METHOD	Hermetically scaled blister pack	Hermetteally, scaled blister pack	Hermetically sealed blister pack

### 6.- PHYSICOCHEMICAL STUDIES

The physical, optical and chemical properties of the lenses as assessed by various test methods show substantial equivalency with the predicate devices as illustrated in the preceding table. Studies were also conducted to verify that leachable substances were either low or below measurable levels to assuage any concerns for its intended use.

### 7.- TOXICOLOGY STUDIES

Sauflon Clariti (somofileon A) Soft (hydrophilic) Contact Lenses with UV Blocker and Handling Tint was assessed using ISO 10993 standards for cytotoxicity, ocular irritation and systemic toxicity. All results passed with no evidence of adverse clinical effects caused by the lens.

### 8.- HUMÁN CLINICAL STUDIES

A clinical study was conducted to evaluate the safety and efficacy of Sauflon Clariti (somofilcon A) Soft (hydrophilic) Contact Lens with UV Blocker and Handling Tint by comparison with Air Optix Aqua hydrophilic contact lenses (Ciba Vision Inc.). Subjects used OptiFree Replenish solution (Alcon Laboratories Inc.) for daily lens maintenance, care and storage. The results of this study showed the safety, acceptability and substantial equivalence of the Sauflon Clariti (somofilcon A) Soft (hydrophilic) Contact Lens with UV Blocker and Handling Tint to the predicate device for its intended use.

### 9.- CONCLUSIONS

Based on the above evaluations the Sauflon Clariti 1 day (somofile on A) Soft (hydrophilic) Daily Disposable Contact Lens with UV Blocker and Handling Tint is substantially equivalent to the predicate, marketed lenses. Based on these evaluations the Sauflon Clariti 1 day (somofile on A) Soft (hydrophilic) Daily Disposable Contact Lens with UV Blocker and Handling Tint has been shown to be safe and effective for its intended use.



March 4, 2014

Food and Drug Administration 10903 New Hampshire Avenue Document Control Center - WO66-G609 Silver Spring, MD 20993-0002

Sauflon Pharmaceuticals Ltd % Dr. Christopher Smejkal Head of Regulatory and Technical Affairs 49-53 York Street, Twickenham Middlesex TW1 3LP United Kingdom

Re: K133472

Trade/Device Name: Sauflon Clariti 1 Day (somofilcon A) Soft (hydrophilic) Daily

Disposable Contact Lens with UV Blocker and Handling Tint

Regulation Number: 21 CFR 886.5925

Regulation Name: Soft (hydrophilic) contact lens

Regulatory Class: Class II Product Code: LPL, MVN Dated: January 22, 2014 Received: January 30, 2014

### Dear Dr. Smejkal:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the <u>Federal Register</u>.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies.

# Page 2 - Dr. Christopher Smejkal

You must comply with all the Act's requirements, including, but not limited to: Registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please contact the Division of Small Manufacturers, International and Consumer Assistance at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address <a href="http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm">http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm</a>. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to <a href="http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm">http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm</a> for the CDRH's Office

of Surveillance and Biometrics/Division of Postmarket Surveillance.

You may obtain other general information on your responsibilities under the Act from the Division of Small Manufacturers, International and Consumer Assistance at its toll-free number

(800) 638-2041 or (301) 796-7100 or at its Internet address http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm.

Sincerely yours,

# Kesia Y. Alexander -S

for Malvina B. Eydelman, M.D.

Director

Division of Ophthalmic and Ear,

Nose and Throat Devices

Office of Device Evaluation

Center for Devices and Radiological Health

Enclosure

### INDICATIONS FOR USE

510(k) Number (if known): K133472

Sauston Clariti 1 Day (somofilcon A) Soft (hydrophilic) Daily Device Name:

Disposable Contact Lens with UV blocker and Handling Tint.

The Sauflon Clariti 1 Day (somofilcon A) Soft (hydrophilic) Indication for use:

Daily Disposable Contact Lens with handling tint is indicated

The SAUFLON CLARITI 1 DAY (somefileon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV blocker and Handling Tint is indicated for daily wear single use only for the correction of refractive ametropia (myopia and hyperopia) in phakic or aphakic persons with non-diseased eyes that may exhibit astigmatism up to 2.00 Diopters that does not interfere with visual acuity.

The SAUFLON CLARITI 1 DAY TORIC (somofileon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV blocker and Handling Tint is indicated for daily wear single use only for the optical correction of refractive ametropia (myopia and hyperopia) in phakic or aphakic persons with non-diseased eyes that may exhibit astigmatism up to 10.00 Diopters.

The SAUFLON CLARITI I DAY MULTIFOCAL (somofileon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV blocker and Handling Tint is indicated for daily wear single use only for the optical correction of refractive ametropia (myopia and hyperopia) and/or presbyopia in phakic or aphakic persons with non-diseased eyes that may require a reading addition of +4.00 Diopters or less and may exhibit astigmatism up to 1.50 Diopters or less.

The SAUFLON CLARITI 1 DAY MULTIFOCAL TORIC (somofileon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV blocker is indicated for daily wear single use only for the optical correction of refractive ametropia (myopia and hyperopia) and/or presbyopia in phakic or aphakic persons with non-diseased eyes that may exhibit astigmatism up to 10.00 Diopters and require a reading addition of +4.00 Diopters or less.

Sauflon Clariti | Day (somofilcon A) Soft (hydrophilic) Daily Disposable Contact lens with UV blocker and Handling Tint help protect against transmission of harmful UV radiation to the comea and into the eye.

Prescription Use

AND/OR

Over-The-Counter Use

(Part 21 CFR 801 Subpart D)

(21 CFR 801 Subpart C)

(PLEASE DO NOT WRITE BELOW THIS LINE - CONTINUE ON ANOTHER PAGE IF NEEDED)

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